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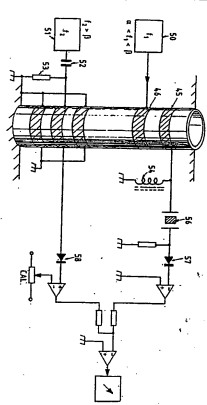
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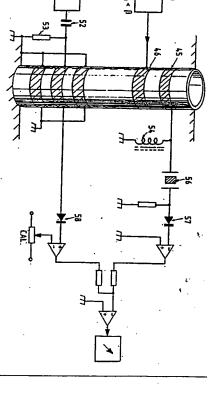
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(54) Tiue: APPARATUS FOR DETERMINING THE PHYSICAL AND/OR CHEMICAL PROPERTIES OF A SAMPLE, PARTICULARLY OF BLOOD





(57) Abstract

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Apparatus and method for correlating certain physical and chemical properties of blood and other samples by means of remote simultaneous multi-frequency dielectric measurement or when one said frequency is applied and compared with an externally entered paramater proportional to the magnitude of a chosen dielectric parameter at another frequency(ies). Apparatus particularly useful for the near instantaneous assessment of the expected sedimentation condition of red blood cells or other fibrinogen and crythrocyte related parameters.

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APPARATUS FOR DETERMINING THE PHYSICAL AND/OR CHEMICAL PROPERTIES OF A SAMPLE, PARTICULARLY OF BLOOD.

5 This invention relates to a non-contacting apparatus parameters and for use with similar evaluations in other and method for invesrigating certain properties of Throughout this embodiment , the term " non-contacting" implies biological media and for more general use with other samples content , sedimentation rate and related physical and chemical blood such as red cell count, haemoglobin and fibrinogen electronic display devices. condition of red cells in blood and certain plasma properties. fibrinogen to establish instantly the expected sedimentation In one example the invention is concerned with measurement of only being limited by the speed of electron flow in circuitry real time electronic calculation and the operation time of means remote from the sample and the terms "instant" and instantaneous" imply very near instant with process times

Protein concentration in biological media is usually assessed by biochemical methods or by methods of physical chemistry such as viscosity measurement and optical rotational dichorism.

Also possible are various forms of spectroscopic analysis and chromatography. In one specific situation, that of whole blood, the proteins with the highest concentration are haemoglobin, found in the erythrocyte nuclei and secondly fibrinogen, found dissolved in the plasma. Fibrinogen concentration is medically important, in that in excess it is a non-specific indicator important in a person. Fibrinogen levels manifest their effects in a variety of different ways; firstly, they effect the

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sedimentation rate of the red cells (erythrocytes) giving "ise to the so called erythrocyte sedimentation rate or e.s.r. and secondly, they cause effect upon the physical and chemical properties of the plasma.

15 slowly under the control of gravity and internal viscoelastic 0 popular. Manifestations of increased fibrinogen levels have traditionally been monitored in pathology laboratories or p.v., more recently a third biochemical assay, the so by two tests, namely; the e.s.r. and the plasma viscosity with clinicians the World over. The e.s.r. test traditionally called c-reactive protein or c.r.p. test has also become more separates from the clearer plamsa fraction and sediments perform , during which time the red cell fraction (haemstocrit) uses about 5 milli-litres of venous blood and takes one hour to chemically separated , there is always the chance , albeit preservative, this is very time -consuming. P.v. and c.r.p. are also time consuming and because in these latter two tests bacterial biohazard remote, that the operatives might become exposed to viral or and white blood fractions have to be physically or E.s.r. tests are however still the most popular a capillary tube or a vacutainer containing

Other blood tests such as cell counting and sizing are also carried out in pathology laboratories using very expensive 25 automated equipment, which needs to sample small quantities of blood in close contact by sucking it through a needle type probe inserted by the equipment into a closed vacutainer. Such cell counters, sometimes referred to as haematological analysers, culter or similar, are extremely sopnisticated

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10 are perhaps the red cell concentration (r.b.c.), the mean gale. These machines yield a myriad of parameters, upto 23 gradients and voltage pulses across individual red or white useful then of the Nevertheless they are non-portable and extremely expensive and in some cases, about the state of nearly all the blood components. blood cells which have been located by electric or hydrodynamic parameters are considered very useful by many physicians in cell volume (m.c.v.) and the haemoglobin content (Hb). These limited by sample throughput and cleansing procedures. Three and operate by application of non-linear electrical field portability, for use for example in G.P. offices, in the be provided by a simpler, cheaper, and general "state of health" assessments. addition to the e.s.r. value in order to make first diagnoses focousing in a narrow ,micron sized, orifice or counting/sizing most important parameters outputted by cell counters technology of greater by the present inventor if such parameters could haematology or hamatological It is considered

It is thus an object then of one aspect of this invention to provide a means to instantly assess blood fibrinogen levels and their related unemical and physical manifestations remotely and to be beable to monitor, also remotely.

is not instant

expensive and because a chemical reaction is involved there is a waiting time before the result is achieved, i.e the output

the problem of haemoglobin has been addressed by previous

Lield, or with Third World applications. Of these parameters

inventors using optical technology and biochemical lysis of

the erythrocytes , however such technology is still quite

any or all of the common red cell parameters referred to above, to help preclude biohazard and to provide an analogue or digital readout of all or any of these parameters. in devices that may or may not be configured as a simple form of haematological analyser, and an instant non-optical, non-cell counter means to determine m.c.v., and/or r.b.c and/or

Haemoglobin content of blood.

10 or have units which are effectively dimensionless but whose 20 (instant sedimentation rate), but also accounting for and numerical dynamic range scales and correlates according an output which can be calibrated in units of In the case of fibrinogen, it is also an object to chosen according to the preferance of the physician etc. which the present inventor chooses to refer to as i.s.r. assessment referred to above ,or according to a new parameter 20 minutes. of e.s.r. , these are not instant but they do the rate are also possible to increase shear forces on the erythrocytes thereby speeding reduce the time required for a measurement down to circa Automated optical systems have been tried for the assessment any or either of the three common methods of fibrinogen Methods where the e.s.r. tube is spinning in order concentration, however provide

Dielectric methods have also been suggested for
the study of time -dependent erythrocyte sedimentation ,
the study of time -dependent erythrocyte sedimentation ,
25 GB 1574681 (Labora Mannheim). In fact , one purpose of this
present embodiment is to provide dielectric
systems that advantageously and differently however provide for
the systems that advantageously and differently however provide for
systems that advantageously and differently however provide for
the systems that advantageously and differently however provide for
systems that advantageously and differently however provide for
the study of time -dependent assessment of e.s.r.

- 10 electrodes in direct contact with the sample, either tubular. GB 1599241 , where using flowing blood in an insulin/glucose control loop , a 500Hz haematocrit level electrode was formed present embodiment to fully distinguish the prior art from in blood , it is considered very important at this stage in have described apparatus for making single frequency dielectric that of this present embodiment. For instance some inventors /or conductivity measurements on liquids using two or more
- 15 (Malcolm -Ellis (Liverpool) Ltd;) these were electrically in a fermenter, where four pointed contacting probes were used configured like the standard d.c. four point probe conductivity measurement. This has also been used at a.c. by Kell (US 4965206)
- 20 according to four point probe theory or principles, is that the electrodes actually make re-iterating, a common disadvantage with the above prior art
- 12 effects , and the chance of cross-infection and biohazard. can give rise to the chance of electrode fouling, electrolytic
- Alternatively alternating voltage of continously varying

- above, the time dependent sedimentation rate haematocrit level of blood and in a separate invention, as measurements on a variety of samples including even the has been described in the prior art for making
- liquor ., at d.c. or very low frequency a.c., GB1460892 annular and four or six in number disposed in alcoholic
- This present invention, quite differently , does not function
- physical contact with the liquid under investigation. This
- 30 that swept frequencies could have been utilised for particles, again by means of contacting electrodes, WO85/ 04481 (swept) frequency has been applied to a suspension of biological (Public Heaith Laboratory Service Board). It is possible

- required was considered unacceptably high by the present inventor however the degree of signal processing which would have been some of the aims of the present invention
- 10 some aspects of this present invention by the use of available also dismissed this possibility. A further disadvantage This need for separate calibration is precluded in separate calibration with its cell full of electrolyte in of the above invention (WO85/04481) is that it required the absence of and the fact that truly "instant" results would not have been biological material in some of its aspects
- dielectric measurement and control where a single pair of machine, EP 0 309 085 A2, (Fischer Scientific Co. one such was a "bang /bang" control device for a drip feed electrodes have been used on the outside of insulating tubes, There have also been a few instances in be needed due to environmental and temperature effects. advantageously can offset problems of calibration which may differential modes as a specific possibility, which very simple
- 17 20 a drip feed tube , if the tube became empty or air-locked. purpose, i:e as a warning or on/off control device (f.e.t. circuit), it caused this crystal oscillator to cease oscillating. This type of prior art is adequate for its the feedback loop of a Pierce crystal oscillator the capacitance fell and finding itself arranged in Pittsburg P.A.), where capacitor plates were placed either side of
- but does not have the applications of this solated rapacitor plates used for actual measurement purposes present embodiment. precision or dynamic range for the An example of

fly ash. . GB 2 115 933 A (Kajaani Oy, (finland)). with a frequency applied is in the assessment of coal content of

- of the impeded by the combined capacitive reactance Essentially such a system worked by monitoring an a.c. level
- erythrocyte sedimentation , but in this case were attached to a test- tube to monitor time -dependent Labora Mannheim , where two plate or curved electrodes Similarly, and previously referred to above is the method of plates , the insulating tube and the fly ash
- 10 their effect was in re-tuning a parallel tuned circuit which being thence connected in parallel with a separate capacitor even be substituted for effectively formed the input tank to a voltage controlled λ single inductor wound around the tube could the capacitor plates in that invention

10 the aims of the present invention.

sensitive enough, stable enough, or fast enough to satisfy

- 15 distinction between the dielectric constant of the haematocrit and thence to a voltage controlled oscillator. Such oscillators Labora Mannheim system that there should be a measureable are not considered stable enough for use with this present invention. It was also a pre-requisite of the
- 20 and plasma fractions the case for vertain pathologies at least. It is unfortunate be true in most cases, it is the contention and experience by least in the low megahertz frequency band, this is not always way of experimental observation of the present inventor that at in all cases, Whilst this may
- 25 that Labora Mannheim did not specify the operating range of the v.u.o. employed in their description. The above contention hav possibly explain why their invention does not appear to have been widely exploited as an e.s.r. monitor

5 oscillator (essentially still a v.c.o.), to monitor the flow Telecom), which used a coil around a tube connected to a search some state of the art techniques, these are niether It is clear then that although the prior art indicates of magnetic particles in a carrier material. properties , another example of this is EP 0157496A2 (Northern with the blood could exploit magnetic as well as dielectric . _abora Mannheim indicated that a single inductor not in contact

25 litherto described in the prior art 20 entered into the calculation circuitry to give a satisfactory 15 principles where unlike the prior art and advantageously and apparatus for applying the above said frequencies, not methods of the prior art . result hitherto not instantly available by other provide new kinds of inductive measurement cell and methods if only one such frequency is applied then an external frequencies are simultaneously applied and employed or where single, prefereably non-varying (i:e stable) parameter will be required to be manually or automatically to it, there are provided either preferably two or more measuring cell , instant methods and apparatus for remote measurement on blood and other fluids based on dielectric Thus it is a further object of the present invention to new and more advanced forms of non-contacting It is a further object to

enemical properties of a sample , blood or other, with aspect , apparatus for determining the physical and/or Accordingly then the present invention consists of in one

means for correlating the required physical and/or chemical sample at each of the said frequencies simultaneously and measuring the magnitude of the dielectric properties of the for applying at least one frequency to the sample , means for means for retaining the sample, means remote from the sample

- frequencies or with an alternative parameter proportional the measuring frequencies with that at the other measuring magnitude of a dielectric property of the sample at one of property of the sample from a simultaneous comparison of the
- method involves inserting and retaining samples, blood or cells linked to electronic circuitry through which external parameters can be entered if necessary. Accordingly the invention consists of non-contacting dielectric measurement Furthermore accordingly, the said apparatus of this
- 15 other, in the said apparatus , applying the said frequencies required physical and/or chemical parameters , and measuring the said magnitudes and correlating the said means of internal electronic (calculating) circuitry. Often providing a scaled readable analogue or digital output by
- 20 one , two or four frequencies are applied . In the case of beta dielectric loss maximum, whereas in the case of four all may be on the high side of this beta loss maximum. The and beta dispersions and the other on the high side of the two frequencies , one may be between the dielectric alpha
- 25 method is ideally suited to assessing protein and cellular concentrations in blood and other biofluids but use of the high frequency tail of the beta dispersion. samples is not ruled out. Protein concentration is by its effects on the position and/or magnitude of furthermore

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as a component of some of the measuring cells available to is electrically insulating and may double as a electrode structures spaced lengthwise on a former which the said apparatus there are provided circumferential

- tube with one, both or no ends open, into which by monitoring the voltage on the transmitting electrode, Furthermore and advantageously, measurement is made either separately insulated open or sealed sample tube might be pushed receiving electrode or both, whereas the
- 10 previous inventors have only monitored the voltage on a receiving electrode, or used the capacitance of electrodes to resonate in parallel with an inductor, to tune a v.c.o.

15 frequencies in kilohertz regions, and assessment of mean cell volume in blood is also made with frequencies in the as those referred to above are employed but assessment is of the number density of red cells by measurement at In another aspect of the cells and method, structures low megahertz regions.

electronic circuit which uniquely and advantageously allows temperature compensation and entry of an external parameter above , but a single frequency is used in conjunction with an In another form the present invention employs structures as is bloom, in order to cell counter or optical haemoglobinometer if the sample S. haemoglobin content from another source such as

fibrinogen content, or fibrinogen related parameter(s). In another form , the present -contacting conductivity and /or dielectric constant of a medium cells are used for assessing changes provide a more precise output of apparatus and its

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following, the temporal evolution of the output parameter. bioreaction , biochemical or biotechnological reaction, by undergoing physical or chemical change e.g.; chemical reaction,

a variable crystal oscillator (v.x.o.) by series upon a crystal controls the frequency and amplitude of in the plane of the former and where the coil via its effect replaced by a single coil or inductor wound around and lying In a further form the prasent apparatus is used for any of the uses referred to above but the electrode structures are

Since output frequency of a v.x.o. can be measured by a counter inclusion in the input oscillating tank ,not output load and not feedback, circuit. employed free running types of oscillator and not a v.x.o. inherently far more stable than those of other inventors that Advantageously such a method is

15 very accurately , down to fractions of a Hertz, there is thus concurrent with the increased stability referred to herein an increased precision and sensitvity over other methods.

a coil structure surounds the apparatus' measuring cell In a further aspect of this present invention,

20 and the coil (inductor) has low impedance tap or link into of the sample are mathematically or empirically related to the which power is fed via a coaxial line from an exciter. Any of the invention configured in this way, since the properties the assays and samples mentioned herein may be attempted with

25 voltage standing wave ratio on or without further d.c. amplification reflectometer or v.s.w.r. meter in that line, provided with the coax line as measured by a

25 this context within the scope of the claims of this invention

phase- locked- loop techniques could also be employed in

recovery methods such as radio frequency amplifiers and/or

evenly spaced , two of which are non-resonant input and two of which receive singly and separately yet (transmit) coils, each sending in a separate single frequency, central former is surrounded by four coils or inductors lying Yet a further aspect of this present invention is a two measurement cell, method and device, in which (circumferentially) around it , evenly or not

through the former walls and sample.

simultaneously these original frequencies after passage

20 through the high impedance of the former and sample holder walls 5 10 In yet a further connected from the electrode to earth which resonates with the Those skilled in the art will appreciate that other signal and would thus be virtually undetectable but for this aspect. electrode which consists of a high Q ferrite cored inductor frequencies would suffer very great attenuation after passage electrode self-capacitance, thereby boosting the recovered low frequency signal. method described earlier but where a signal there is operation as per the electrode based two frequency recovery technique is employed on the low frequency receive signals This is aspect of this present invention of kilohertz advantageous because

10 with and may contain any of the aforesaid or following aspects effect of r.b.c., m.c.v., Hb, various other proteins, cell present inventor that these "norms" arise due to the combined of this present embodiment. It is further ascerted by the general health status indicator based on the observation aspect of this invention to provide means of an electronic from which the sample was acquired , thus it is yet one further .1 GHz are related to the general state of health of the individual that there are "norms" of dielectric response at each frequency sample measured at individual frequencies in the range 10KHz extstyle -It is an observation of the present inventor that the dielectric (capacitive and conductive) facets of a pathological blood the radio frequency continuum and that this can be used

15 upon the loss peak maxima magnitudes and positions in in the approximate frequency range $0.5-60\ \mathrm{MHz}$. dispersion of blood , with these dispersive phenomena lying frequency space of the double or multiple dielectric Beta

membrane leakiness and plasma electrolyte strength

25 constant chemical and physical composition and dielectric 20 embodiments consists of cells, methods , means and devices of sample containers, should these vary from container to optics some of the physical dimensions and dielectric properties capable of measuring without contact and without the use of container if said containers are filled with fluid of In yet a further form the invention in any of its previous

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10 connecting them to a differential output stage, this aspect with the sample being placed in /measured by one member of mechanical and electronic , in each of the said sets and then or electroloyte etc; being placed in /measured by second said set and a dummy sample, containing for example air, water using two identical sets of said cells , devices or apparatus this invention may be operated in a differential mode, i:e methods, and apparatus referred to herein as belonging to In a final aspect of this invention, any of the measuring cells will tend to be cancelled by the differential stage. of the present invention allows for the provision of member of said set. improved results as environmental efects such as temperature By employing identical components,

15 are primarily illustrated as device(s) for determining protein and cellular concentration in liquids, preferably whole blood within the scope of the claims. detail shown , since various modifications could be made therein without contact, it is not intended to be limited to the precise Although the invention and all its embodiments described herein

20 The invention and some of the advantages thereof will now be drawings in which; described more fully by way of reference to the accompanying

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Figure 1, illustrates the two frequency measurement cell.
with its insulating former and outer annular (circumferential)
electrode structure, for use in this invention;

figure 2, illustrates the method of monitoring voltage at the transmit electrode in this invention and shows the stray capacitance path to earth, according to this invention;

Figure 3, is a diagram illustrating the method of signal recovery, for boosting kilohertz signals after passage through the former and sample, according to this invention;

10 Figure 4 , is a diagram of the alternative measurement cell with an inductor connected to a variable crystal oscillator, for use with this invention;

Figure 5, is a diagram of the measurement cell, former, tapped coil and device showing manner of connection to 15 voltage standing wavemeter (reflectometer), according

to this invention;

Figure 6, is a diagram of the two frequency four coil measurement cell according to this invention;

Figure 7, illustrates a block diagram of a two frequency method 20 and device for the measurement of protein, preferably fibrinogen in blood, device can also be used to measure red cell concentration and/or mean cell volume by appropriate adjustment

of frequency pairs according to this invention;

Figure 8, illustrates a four frequency method and device for the measurement of protein, preferably fibrinogen in blood, according to this invention;

where a single frequency device is used in conjunction with an external entry parameter to yield a new parameter, where entry parameter is preferably haemoglobin content, to yield fibrinogen content or related parameter, at output 10 if sample is blood, and finally,

Figure 10 , illustrates the differential mode, according to this invention.

Referring to figure 1, the two frequency measurement cell, 10 and 11 are circumferential transmit electrodes remote 15 from the sample, they are usually, although not exclusively fabricated from thin brass shim. Prequencies f, and f, are simultaneously passed into 10 and 11. 12 and 13 are two similar receiving electrodes from which f, and f, are simultaneously recovered. 14 is a central grounded electrode to 20 minimise stray signal leak along the surface of former/tube 15. 16 and 17 are earthed ground-planes to minimise r.f. radiation from the cell.

Referring to rigure 2, the method of measuring voltage at the

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trimmer capacitor , 20 a resistor , usually although not similar stable exciter. 19 is a 10 picofarad (or thereabouts) transmit electrode; 18 is a crystal controlled oscillator or

- relatively high r.f. voltage. This method has the advantage that detection is made at a exclusively in the range 5 -25 k ohms, 21 is a signal diode. blood or similar into the orifice 22, this change occurs impedance at the transmit electrode to a value which is easily influenced or changed by introduction of a sample tube containing 19 and 20 adjust the effective
- 10 due to leak of signal to earth and the impedance at the transmit electrode being too high to sustain constant current sample is introduced. Terminals 23 flow, thus the voltage an electronic voltmeter for interpretation. on this electrode will fall when a are thence connected
- 20 boost the recovered signal appearing for detection at 28. 15 Referring next to figure 3, the method of signal recovery for high Q resonance with ferrite cored inductor 27 in order to the receiving electrode whose self-capacitance 26 brings about presently ,though not exclusively, a 160 kHz sine -wave and 25 is kilohertz frequencies. 24 is a kilohertz frequency generator,

25 in series with crystal 31 to form the input tank circuit of v.x.o.(variable crystal oscillator) 32. Reference to figure 4 shows the alternative measurement cell and amplitude of 32 will differ invention. single frequency v.x.o. method used with this present Coil 30 is wound around former 29 and is connected The output frequency

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simple on/ off bang/bang control. used coil in feedback circuit of crystal oscillators for stability of a v.x.o., and is superior to those which have inventions which have used a v.c.o. due to inherently higher to amplitude and frequency of 32 . Method is superior to those Thus physical and chemical properties of sample may be related any of the sample properties is temporally unstable. when 29 is empty and when 29 contains a sample in its own tube They will also differ from sample to sample and will drift if

- 10 Reference to figure 5, shows continuous wave v.s.w.r. method which is a low impedance tap point or a link. where 33 is an inductor but where an essential feature of resonates with a capacitor , either self capacitance of inductor and former or external additional paraliel 33
- 15 capacitance . Power is fed into 33 from exciter 35 via reflectometer
- d.c. amplification. When sample tube is pushed into orifice or voltage standing wave meter 34. of former, resonant frequency of system alters slightly, 34 may or may not require
- 25 a sample. 20 causing an alteration in the amount of power absorbed by 33 and within this definition is covered temporal instability of v.s.w.r. is sensed and measured by 34, thus the reading reflected back towards 35 , the change in this reflection 34 relates to physical and chemical properties of the sample, 2

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by non-resonant link inductors 37 and 38 respectively. Power is passed in at two frequencies f_1 and f_2 simultaneously Reference to figure b , shows the inductive variant of figure 1 , a two frequency four coil measurement cell used in this invention.

Said frequencies are recovered after passage through the mathematical comparison, calculation or processing then application and sample type. follows on the voltages v, and v, depending on the precise tuned circuits 39/41 and 40/42. former , sample tube and sample by resonant recovery at parallel Any chosen degree of

Thus appropriate mathematical manipulation

to a complex mathematical

function

involving both.

10 Reference to figure 7 shows a specific use of the invention as 58 sperate exactly as in accordance with the equivalent parts in the voltage monitoring system described in figure 2. exclusively around 50 MHz). 48 and 49 , 51 and 52 , 47/49 and tail of the dielectric beta dispersion (usually although not of fibrinogen in blood . a device, in block diagram form, preferably for the measurement Frequency f is on the high frequency

20 and 45, total protein content being mainly haemoglobin and fibrinogen. In the case of blood, the detected voltage is related to the Frequency f lies between the alpha and beta dispersions

25 of high frequency signal which may have strayed into this part a series quartz crystal or similar filter to remove any traces However , 56 is an extra component which takes the form of frequency recovery method described by reference to figure 3. with their equivalent counterparts in the kilohertz The vircuit where it is unwanted. The voltage at the detector 46 , 50 and 54 operate exactly as in accord

Said voltage at 57 is also weakly dependent on haemoglobin of pathological samples (private study of the present inventor). is blood, and this number density in turn correlates to a large 57 is related to the number density of erythrocytes, if sample concentration direct and also on extent with sample haemoglobin content, for the vast majority mean cell volmue according

20 15 inventor chooses to refer to as the i.s.r. (instant sedimentation 10 simplest comprising two operational amplifiers) can remove is no reason why the output should not be scaled in order may be arranged to yield an output parameter which the present protein function, to leave remaining a signal contribution which an approximate contribution due to hasmoglobin from the total of the signals from detectors 57 and 58 in circuit 59 (at in magnitude and dynamic range of the more traditional e.s.r., depends mainly on fibrinogen levels. The output scale factor present claims herein. parameters and indeed this is within the scope of the reading covering the Those skilled in the art however will appreciate that there rate), if the sample is blood, this parameter may be scaled parameter give an "instant " p.v. reading which physicians are more used to interpreting equivalent dynamic ranges of these two or an "instant" c.r.p. its

25 Referring next to figure 8 , this illustrates a block diagram of the four frequency well, measurement method and device for use

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proteins, e.g. haemoglobin and fibrinogen, if the sample is blood, beta dispersion are influenced in different ways by different frequency space) of the high frequency tail of the dielectric with this present invention. Because different parts (in

- 10 order of 30 MHz , and, $f_{f \mu}$ is of the order of 50 MHz. it is possible to obtain an estimate of fibrinogen levels by Frequencies f_t - f_{ψ} are passed in through electrodes 60-63 and order of 17 MHz, $\mathbf{f_2}$ is of the order of 20 MHz , frequency range 15 -60 MHz (usual but not exclusive range within simultaneous four frequency dielectric measurement in scope of this present invention). Usually frequency \mathbf{f}_{\parallel} is of the is of the
- 15 from the 17 MHz filter and detector by that derived for the 20 MHz signal . for fibrinogen, then wieghted subtraction in 75 tends to enhance similar components in repect of Heamoglobin but somewhat different ${
 m f_3/f_4}$. For blood as a sample, output functions of 73 and 74 have 73 is an analogue divider which divides the detected voltage Likewise , 74 performs a similar operation for

centered on f_i - f_{μ} respectively to assist with signal recovery.

out through 65-68 inclusive. 69-72 are narrow band-pass filters

- the effect of fibrinogen and suppress the effect of haemoglobin At this point in the circuit the fibrinogen function is almost offset is provided by 76 so that the output parameter may be displayed at 77. linear but is superimposed on a d.c. level , thus an appropriate Those skilled in the art will appreciate that
- the technique is not limited within the scope of the claims to only blood as a sample and indeed any system containing cellular be amenable to this kind of treatment. When the sample is biomass and protein together or even mixtures of proteins will

of determining fibrinogen but because four frequencies are blood, this aspect of the invention is a most accurate way employed, very careful adjusment and initial calibration

solutions is necessary and temperature compensation of 73-76 is also desirable. initially with pathological samples and latterly with electrolyte technologically challenging. Thus this aspect of the invention is

10 assessment when a numeric entry parameter (e:g haemoglobin) and is used as the said external entry parameter similar cell counter or biochemical optical haemoglobinometer known or available from another source such as Coulter or is available or known. aspect of the invention concerned with fibrinogen or protein Referring next to figure 9 , the block diagram of If Haemoglobin content of blood is

- 15 then the invention configured according to this aspect system 78-81 operate to the drawing , the main component parts of the assessment of fibrinogen level. Referring then can be used to provide a simpler and more accurate in exactly the
- 20 same accord as their equivalent parts indicated in figure and temperature is compensated for using potentiometer 82. Those The digital voltmeter 84 is used with a differential input skilled in the art will appreciate automatic compensation also to be possible within the scope of the present claims. 83
- 25 Haemoglobin entry circuit is also shown for simplicity as a potentiometer, but in reality in the working demonstrator instrument it comprises of a set of rocker or chumbwheel type

entry, both analogue and digital within the scope of the claims appreciate that there are several other means of haemoglobin value to the nearest whole unit. Those skilled in the art will switches and it is usually adequate to enter the Haemoglobin

- of this present invention, including for example; acquistion of the haemoglobin level by direct connection
- protein content and the differential action of 84 removes from because the voltage at 81 is an inverse function of the total according to this aspect addition of the temperature compensation voltage. Those skilled this the hasmoglobin contribution and simultaneously allows haemoglobinometer. to the electronic the art will appreciate that The action of the system is achieved circuitry of a cell counter could be used with the invention configured
- within the scope of the claims of this invention, and that if to yield a heamoglobin value at its output, haemoglobin that the system could be configured "in reverse" manually acquired e.s.r. value were available instead of

multicomponent fluid systems other than blood

- 20 within the scope of these present claims. Those skilled in applied through just one electrode or inductor , within the art will appreciate that simultaneous frequencies may be combiners and/or directional coupling techniques scope of the claims of this present invention by using power the
- own container, said container being a tube, vacutainer, referred to in this present embodiment and by way of reference employing any of the said cells ,means, methods and devices Another feature which should not be overlooked when the drawings is that when the sample is contained in
- capillary etc, with open or sealed end(s), aforesaid container

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15 invention referred to above and in the claims herein , 10 It will however be appreciated by those skilled in the art dimensions vary (from container to container) , particularly Such errors arise from variations in the air gap capacitance should be a snug push fit into former /tube of said cell etc; namely that if samples of fixed chemical and dielectric be turned on its head to yield yet a futher aspect of the Furthermore they will appreciate that this problem may where the air gap is that between the said container and former measurement produced by methods and devices herein may arise. air gap (although not all the air is displaced) between this figures 1-9, and there should not be excessive slack or excess property are employed in sample containers of nominally the automatically by tube size correction techniques. that such errors can be reduced/corrected for manually or the internal and external diameters, then errors in the container and the former inner walls. property, then said cells , methods , means and devices may same size but with slight variations in size or dielectric If the container

formers as illustrated in any of the prior drawings in this Referring finally to figure 10. 85 is the sample tube and circuits associated with any of means , methods and devices present embodiment. 87 is the dummy or control sample tube. 86 & 88 are identical 89 & 90 are identical electronic

be used to measure a physical dimension of sample container

without the use of a rule, calipers , micrometer other gauges

1.07

in this invention. 41 is a difference amplifier and 92 an appropriately scaled output device/ display. Effects of temperature and other environmental factors tend to be cancelled by this arrangemnet, thus making the invention according to this aspect more stable and accurate than those previous disclosures which do not employ a differential mode.

10 open ended sample tube. Nothing in this embodiment prevents been considered to be on the whole stationary in κ closed or wise retained, but yet with operation in accordance with the aforesaid formers could be fabricated in a "turned inside appreciate that it is hereby also disclosed that the new methods means devices etc. herein can also be made to work when there ohvious advantages , however those skilled in the art will exclusively to non-contacting systems mainly to emphasise their claims of this present invention. of on the inside and with their ends closed to prevent fluid out" manner i:e with their electrodes or inductors disposed this present embodiment, reference has been made so far entry or contact with said electrodes or inductors, thus will be appreciated by those skilled in the art, that the Throughout this embodiment, the sample by way of example has is contact with the sample material by minimal modification. forming probes which could then be dipped into samples otherthe sample from being a flowing or moving sample, in which case formers referred to in every aspect herein would be of the variety with both ends open. Furthermore, it Furthermore, throughout

the said cells .methods ,means and devices referred to herein may be provided with manual or automatic means of sample mixing handling, labelling etc; and results, analogue or digital, could also be computer stored or on a print-out, and samples may or 5 may not be aspirated from their original containers into

Furthermore, nothing in this present invention prevents the sample from being biomaterial in vivo, small e.g. cells or large e.g. human body digits, limbs etc.

second or subsequent containers.

Furthermore those skilled in the art will appreciate that there is scope for modification in the aspects of the embodiment that refer to simultaneous multi-frequency excitation and reception 10 since digital as well as analogue methods can be used here and pseudo instantaneous output may be obtained by using fast frequency steps or sweeps of frequencies applied to transmit electrodes. Furthermore in all aspects where diode detection is employed within this present embodiment, see particularly

figures 2 and 3 and figures 6-9, this can be replaced by phase

sensitive detection as a viable alternative with the dual

consequence of added sensitivity and two component information

from the real and imaginary part analysis, advantageous since in reality samples exhibit complex dielectric behaviour and

20 dielectric constant, sometimes referred to as permitivity has such real and imaginary parts. For a said sample dielectric property the present inventor states the real part of permitivity is a measure of the sample a.c. capacitance and with the present invention the apparatus using circumferential electrodes will respond mainly to this capacitive facet, whereas that using coils will respond more strongin to the imaginary part of the permitivity (loss) or conductive facet.

furthermore, those skirled in the art will appreciate that all

CLAINS

- Apparatus for determining the physical and/ or chemical properties of a sample, comprising means for retaining the sample of a sample, comprising means for retaining the sample, means remote from the sample for applying at least one frequency to the sample, means for measuring magnitude of dielectric properties of the sample at each of said frequencies simultaneously, and means for correlating required physical and /or chemical property of the sample from a simultaneous comparison of the magnitude of said dielectric property at one of the measured frequencies with that at the other measured frequencies or with an alternative parameter proportional thereto.
- Apparatus according to claim 1, where said frequencies are stable and non-varying.
- Apparatus according to claim 1, where said frequencies are applied through circumferential electrodes spaced in line.
- Apparatus according to claim 1, where said frequencies are received after passage through sample by circumferential electrodes.
- Apparatus according to claim 1, where said frequencies are applied through inductors.

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- b. Apparatus according to claim 1. where said frequencies are applied by link coupled inductor(s) connected to an exciter via a voltage standing wave meter (reflectometer).
- Apparatus according to claim 1, where said frequencies are applied through tapped inductors.
- Apparatus according to claim 1, where said frequencies are applied by a variable crystal oscillator (v.x.o.).
- Apparatus according to claim 1, where said frequencies are received at parallel resonance.
- 10. Apparatus according to claim 1, where said frequencies are received as reflected power in a low impedance line, after reflection from link coupled inductor.
- 11. Apparatus according to claim 1, where said frequencies are received as reflected power in a low impedance line, after reflection from a tap coupled inductor.
- 12. Apparatus according to claim 1, where means for measuring magnitude of dielectric properties involves r.f voltage measurement and/or d.c voltage measurement after detection, said voltage arising from receive electrodes and for inductors.

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- 14. Apparatus according to claim 1 and 8, where means for measuring magnitude of dielectric properties involves monitoring output frequency of said v.x.o.
- 15. Apparatus according to claim 1 and 8, where means for measuring magnitude of dielectric properties involves monitoring amplitude output of said v.x.o.
- 16. Apparatus according to claims 1, 6 and 10, where means for monitoring magnitude of dielectric properties involves measuring voltage standing wave ratio in feed line to link.
- 17. Apparatus according to claims 1, 7 and 11, where means for monitoring magnitude of dielectric properties involves measuring voltage standing wave ratio in feed line to tap.
- 18. Apparatus according to claim 1, where said magnitude is mainly that of capacitive facet of dielectric property at each simultaneous measurement frequency.
- 19. Apparatus according to claim 1, where said magnitude is mainly of conductive facet of dielectric property at each simultaneous measurement frequency.

- 20. Apparatus according to claim 1, where said magnitude of dielectric properties is dependent on both capacitive and conductive facets of sample.
- 21. Apparatus according to claim 1, wherein said means for correlating physical and/or chemical properties of sample from simultaneous magnitude of comparison of dielectric property at one measured frequency with that at other(s) is achieved by internal electronic circuit.
- 22. Apparatus according to claim 1, where correlation of said properties from said simultaneous comparison of magnitude of dielectric property at one measured frequency with alternative parameter proportional to said magnitude at other frequencies employs an electronic cicuit for manual entry of said alternative parameter.
- 23. Apparatus according to claim 1 and claim 22, as per claim 22, except where said electronic circuit permits the automatic entry of said alternative parameter from an external source e.g. cell counter and/or haemoglobinometer.
- appparatus as in claim 1, operated in a differential mode.

- 25. Method for determining the physical and /or chemical 1, comprising the steps of; inserting /retaining the parameter which is correlate of required physical or frequencies or with alternative parameter proportional with that at other (simultaneous) measurement dielectric property as at one of applied frequencies chosen physical and/or chemical property of sample sample, applying at least one frequency, correlating chemical property. thereto, then scaling and reading instantaneous output from simultaneous comparison of magnitude of desired properties of a sample employing apparatus as in claim
- 26. Method according to claim 25 above, wherein the sample
- 27. Method according to claim 25, wherein the sample is any blood fraction and /or component.
- 28. Nethod according to claims 25 and 26, wherein the P.v. , c-reactive protein and i.s.r. or all of fibrinogen related parameters namely; e.s.r., haemoglobin contnet (Hb); fibrinogen content or any correlate parameter is any or all of the following: red cell count (r.b.c.); mean cell volume (m.c.v.);
- 29. Method according to claim 25, where the sample is biofluid other than blood,

- 30 . Method according to claim 25, where the sample is a liquid other than biofluid.
- 31. Method according to claim 25 and 26 but where the output is scaled in arbitary units of "general health status indication"
- 31. Method according to claim 25 where the sample is a chemical and dielectric property and an insulating becomes a physical dimnsion of tube/holder. tube /holder where thus said required correlate composite comprising a liquid of constant physical
- 32. Method according to claim 25 where the sample, is a digit of the human body.
- 33. Method according to claim 25 where the sample is a limb of the human body.

<u>.</u>33

WIENDED CLAIMS
[received by the International Bureau on 23 July 1993 (23.07.93);
original claim 1 amended; claims 25-34 replaced by amended claims 29-38
new claims 2-5, and 39-45 added; claims 2-24 renumbered as claims 6-28
wherein claims 9,13,16,18,19 and 22-26 are amended (7 pages)]

- magnitudes of dielectric properties of sample arising two or more parameters comprise of from two or said physical and/or chemical for retaining the sample, and comprising a chosen Apparatus for property(ies) at alternative measurement frequency (ies). magnitude is proportional to the sample dielectric frequencies and /or an externally entered parameter whose from simultaneous measurement measuring cell chemical properties of a sample , comprising means more parameters , where said determination of the and comprising means of correlating properties of sample at various measurement physical and/
- ٣ Apparatus as in claim 1 wherein the said determination is
- Apparatus as in claim 1 wherein the said determination is contactless, ise without direct electrical contact to the
- are in the range Apparatus as in claim 1 , where the said various frequencies above 10 KHz and below 1000 MHz
- ŗ Apparatus as simultaneously with frequency application and dielectric property measurement. in claim 1 where the said correlation occurs

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- Apparatus according to claim 1, where said frequencies are stable and non-varying.
- Apparatus according to claim 1, where said frequencies are applied through circumferential electrodes spaced in line.
- Apparatus according to claim 1, where said simultaneous frequencies are received after passage through sample by circumferential electrodes
- Apparatus according to claim 1, simultaneous frequencies are applied through multiple inductors. where said
- 10. Apparatus according to claim 1, where said frequencies an exciter via a voltage standing wave meter are applied by link coupled inductor(s) connected (reflectometer). ť
- Apparatus according to claim 1, where said frequencies are applied through tapped inductors.
- 12. Apparatus according to claim 1, are applied by variable crystal oscillator (v.x.o.). where said frequencies
- are received at parallel resonance . Apparatus according to claim 1, where said frequencies

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- 15. Apparatus according to claim 1, where said frequencies line, after reflection from a tap coupled inductor. are received as reflected power in a low impedance
- electrodes and /or inductors, after frequency selective after detection, said voltage arising from receive voltage measurement and/or d.c voltage measurement frequency involves simultaneous r.f. magnitude of dielectric properties at each simultaneous Apparatus according to claim 1, where means for measuring (filtered) recovery.
- 17. Apparatus according to claim 1, where means for measuring magnitude of dielectric properties involves measurement of voltage levels at transmit electrodes.
- 18. Apparatus according to claim 1, where means for measuring magnitude of dielectric properties involves monitoring output frequency of said v.x.o.
- 19. measuring magnitude of dielectric properties Apparatus according to claim 1 , where means for involves monitoring amplitude output of said v.x.o.

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- 20. Apparatus according to claims 1, 10 and 14, line to link. involves measuring voltage standing wave ratio in feed for monitoring magnitude of dielectric properties where means
- Apparatus according to claims 1, 11 and 15, where means feed line to tap. involves measuring voltage standing wave ratio in for monitoring magnitude of dielectric properties
- 22. Apparatus according to claim 1, where said magnitudes of said dielectric properties ares mainly those of property as at each simultaneous measurement frequency. capacitive facet of the dielectric
- Apparatus according to claim 1, conductive facet of the dielectric property as at said dielectric properties are mainly those of the each simultaneous measurement frequency. where said magnitudes of
- 24. and conductive facets of sample. said dielectric properties are dependent on both capacitive Apparatus according to claim 1, where said magnitudes of
- 25. correlating physical and/or chemical properties of Apparatus according to claim 1, wherein said means for circuit. dielectric property at one of simultaneous frequencies with sample from said magnitude of comparison of that at other(s) is achieved by internal electronic

- 26. Apparatus according to claim 1, where correlation of alternative parameter. frequency with alternative external entry parameter properties from said simultaneous comparison of said physical and/or chemical sample employs an electronic cicuit for manual entry of said magnitude of dielectric property at one measured
- 27. Apparatus according to claim 1 and claim 26, as per claim 26 , except where said electronic circuit permits the haemoglobinometer external source e.g. cell counter and/or automatic entry of said alternative parameter from an
- 28. Appparatus as in claim 1, operated in a differential mode.
- 29. Apparatus according to claim 1 where the said preferably sample dielectric Beta dispersion frequency band. two or more frequencies may be close to or within the
- 30. Apparatus according to claim 1 above, capable of use with
- 31. Apparatus according to claim 1, capable of use with any blood fraction and /or component.

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- 32. Apparatus according to claims 1 and 30, wherein the correlate parameter is any or all of the following: erythrocyte sedimentation rate). c-reactive protein and i.s.r. (instantly predicted or all of fibrinogen related parameters namely; p.v haemoglobin content (Hb) ; fibrinogen content or any red cell count (r.b.c.); mean cell volume (m.c.v.);
- 33. Apparatus according to claim 1 capable of use with biofluid other than blood.
- 34. Apparatus according to claim 1 capable of use with liquid other than biofluid.
- Apparatus according to claims 1 and 30 but where the output is scaled in arbitary units of "general health status indication".
- 36. Apparatus according to claim 1 capable of using a sample chemical and dielectric property and an insulating composite comprising a liquid of constant physical, becomes a physical dimension of the tube and /or holder tube /holder where thus the said correlate
- 37. Apparatus according to claim 1 where the sample is a digit of the human body.
- 38. Appartus according to claim 1 where the sample is a limb of the human body.

- 39. Apparatus as in claim 1, but where the frequencies are applied as fast frequency steps or sweeps rather than simultaneously and thus wherein said correlation output will suffer a slight time delay and hence be described as psuedo instantaneous.
- 40. Apparatus as in claim 1 , but where the said frequencies are all applied through a single electrode.
- 41. Apparatus as in claim 1 , but where the said frequencies are all applied through a single inductor.
- 42. Apparatus as in claim 1 capable of use with a sample where protein concentration is assessed by its effects on the position and magnitude of the high frequency tail of the beta dispersion.
- 43. Apparatus as in claim 1 where digital and analogue methods are employed.
- 44. Apparatus as in claim 1 wherein phase sensitive detection is employed.
- 45. Apparatus as in claim 1 but wherein the said chosen measuring cell is fabricated as a probe.

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STATEMENT UNDER ARTICLE 19

Biology Society, vol. 10, November 1988, New Orleans, pp 761-762 of the Annual Conference of the IKEK Engineering in Medicine and where that sample is respectively blood and a human finger and the of the International Search Report where contact is made to a sample sample has been emphasised, to distinguish it from those citations i:e, use of this invention without direct electrical contact to the not in concept contained in the full present description. changes and amendments do not draw upon anything which is claims 2-5, ought now to distinguish this present invention made to claim 1 which it is felt, together with the new 39 - 45 have been added. Various claims have been amended, extra claims 2-5 and cells and unlike the current invention , have antennas for remote distinguish it from cited apparatus which does not use measuring has an integral measuring cell as part of the apparatus thus to Importantly, claim 1 has been amended to show that this invention respective citations concerned are : FR, A, 2 201 762 and Proceedings is highlighted as is , the "contactless nature", in new claim 3, In the new claim 2, the "instant" nature of the determination more fully from those of the prior art. The may correlate the sample properties from dielectric properties US, ,A, 3 483 860 as have also been cited in the International to the number of applied frequencies and input /output parameters aspects of this present invention have been clarified with regard Search report. investigation of biological targets such as : US,A, 4 135 131 and implication from lines 5-13 of the new claim 1 that the invention required for the correlation of sample properties to show by Furthermore in the amended claim 1 the technical Most importantly, changes have been

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of erythrocyte sedimentation rate, namely WO , A, assessment (as opposed to instant as in this present invention) substantially lower frequencies to make a time dependent US,A, 4 257 001, and to distinguish it from the one which uses GB, A, 1 084 860; DE, A, 3 722 213; DE, A, 3 637 549 and citations are namely: US, A, 4 135 131; US , A,3 483 860; than those of the present invention e:g, those cited which employ microwaves in the range 1-100 GHz applied either by cavities, antennas or waveguides as single or swept frequencies , those and different hardware for the application of those frequencies International Search Report which employ different frequencies to more fully distinguish this invention from those of the see also pages 10,13,17 and 21 of the existing description. has been included in the alternative claim 4, Finally the working frequency range of the present invention clarified , establish sufficent difference to the prior art. do not of course have the added advantage of the said externally FR, A, 2 378 282 and EP , A, 0 157 496 entered parameter and thus it is hoped these differences, now which

Further minor amendments have been made to various other of the claims only drawing on material from the body of the description and not significantly altering the understanding therein. All reference to a method has been removed from original claims 25-34, now replaced by amended claims numbered 29-38. Additional claims 39-45 have been added.

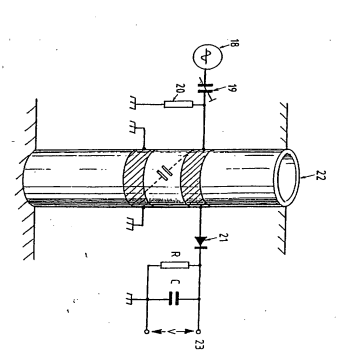
this was a condensed way of making the same statement but obviously it was phrased in such a way that the Search unearthed several single frequency resonance Q techniques namely: GB, A, 2 130 728; GB,A, 595 720; GB, A, 2 248 301;

wording of

gure 9. It was assumed that the original claim 1 carried the same meaning , ise that

-25 and figure 9.

dependent on the application of multiple measurement frequencies two or more in number , see also page 8 , line 16 of the original description and/or by the use of a single frequency or one or more frequencies together with the use of the said externally entered parameter , see page 8 lines 18-22 , page 10, lines 18



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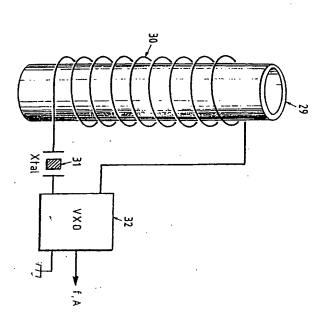
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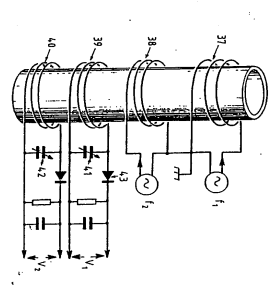
Fig.3.



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TAP جے capacitor کے

Fig . S.



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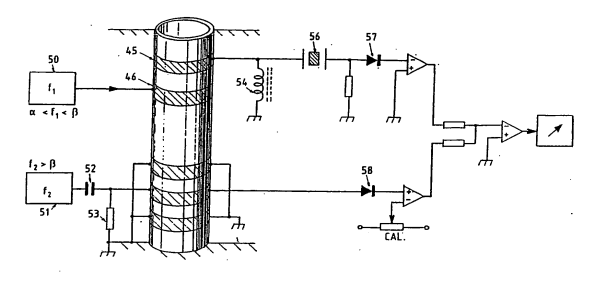


Fig. 7 .

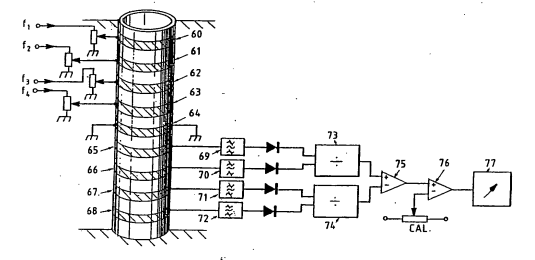
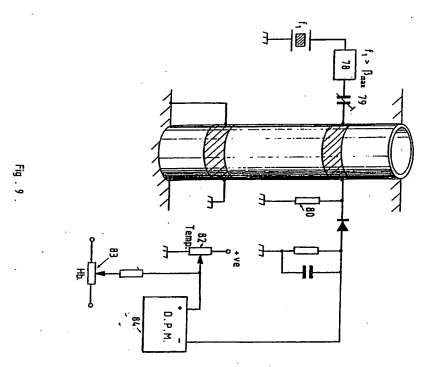
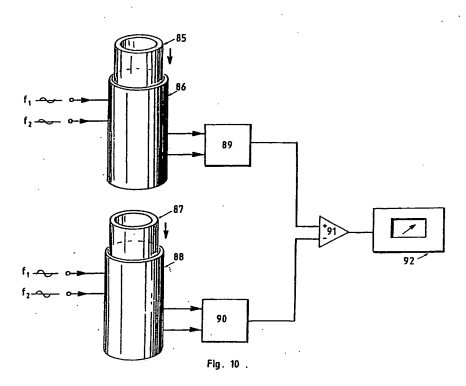


Fig . 8 .

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INTERNATIONAL SEARCH REPORT

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17 March 1981 see claims 1-34 WO,A,9 109 295 27 June 1991 see claims 1-15 DE,A,3 722 213 12 January 198 see column 1, 1 GB,A,2 248 301 1 April 1992 see page 18 EP,A,0 157 496 9 October 1985 FR,A,2 378 282 18 August 1978 see claims 1-4 DE,A,3 637 549 11 May 1988 see column 1 - US,A,3 483 860 16 December 15 see claims 1-3 PROCEEDINGS OF CONFERENCE OF 1 MEDICINE AND EVOL. 10, Novembroom 10, Novembroom 11, Novembroom 12, Novembroom 12, Novembroom 13, Novembroom 14, Novembroom 15, Novembroom 16, Novembroom 16, Novembroom 17, Novembroom 17, Novembroom 18, Novembroom 19, Novem	17 March 1981 see claims 1-34 W0,A,9 109 295 27 June 1991 see claims 1-15 DE,A,3 722 213 12 January 196 see column 1, 1 April 1992 see page 18 EP,A,0 157 496 9 October 1985 FR,A,2 378 282 18 August 1978 see claims 1-4 DE,A,3 637 549 11 May 1988 see column 1 - US,A,3 483 860 16 December 15 see claims 1-3 PROCEEDINGS OF CONFERENCE OF 1 MEDICINE AND E vol. 10, Novembrages 761 - 76 J.M. MCKEE ET 76	III. DOCUMENTS CONSIDERED TO BE RELEVANT CATEGOD® CIRCOD® CIRCOD
WO, A, 9 109 295 27 June 1991 see claims 1-15 DE, A, 3 722 213 12 January 198 see column 1, 1 GB, A, 2 248 301 1 April 1992 see page 18 EP, A, 0 157 496 9 October 1985 FR, A, 2 378 282 18 August 1978 see claims 1-4 DE, A, 3 637 549 11 May 1988 see column 1 - US, A, 3 483 860 16 December 15 see claims 1-3 PROCEEDINGS OF CONFERENCE OF 1 MEDICINE AND WIEDLICINE AND Vol. 10, Novembrages 761 - 766 J.M. MCKEE ET J IMPEDANCE MEASI see page 761 - 10 US, A, 4 135 131 16 January 199	WO,A,9 1 27 June see clat DE,A,3 7 12 Janua see colu GB,A,2 2 1 April see page EP,A,0 1 9 Octobe FR,A,2 3 18 Augus see clat ODE,A,3 6 11 May 1 15 Decan 16 Decan PROCEEDI CONFEREN MEDICINE VOl. 10, pages 76 J.M. MEDICINE INPEDANO see page US,A,4 1 16 Janua	×
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US,A,4 135 16 January	US,A,4 135 16 January	
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ANNEX TO THE INTERNATIONAL SEARCH REPORT
ON INTERNATIONAL PATENT APPLICATION NO.
SA 70982

The members are as contained in the European Princist Oliber ELFF the on The European Patent Office is in no way liable for these particulars which are merely given for the purpose of information.	This among lists the patent family members relating to the patent documents cited in the above-mentioned international search report
·n 18/06/93) report

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	us-A-	None	None	DE-A- SE-A- US-A-	JP-A-	None	None.	SE-B-	None	None	AU-8-	NL-A-	None	None	Pater
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